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## CODING SEQUENCES OF THE HUMAN BRCA1 GENE

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### FIELD OF THE INVENTION

This invention relates to a gene which has been associated with breast and ovarian cancer where the gene is found to be mutated. More specifically, this invention relates to the three coding sequences of the BRCA1 gene BRCA1(omi1),  
10 BRCA1(omi2), and BRCA1(omi3)) isolated from human subjects.

### BACKGROUND OF THE INVENTION

It has been estimated that about 5-10% of breast cancer is inherited Rowell, S., et al., *American Journal of Human Genetics* 55:861-865 (1994). Located on  
15 chromosome 17, BRCA1 is the first gene identified to be conferring increased risk for breast and ovarian cancer. Miki et al., *Science* 266:66-71 (1994). Mutations in this "tumor suppressor" gene are thought to account for roughly 45% of inherited breast cancer and 80-90% of families with increased risk of early onset breast and ovarian cancer. Easton et al., *American Journal of Human Genetics*  
20 52:678-701 (1993).

Locating one or more mutations in the BRCA1 region of chromosome 17 provides a promising approach to reducing the high incidence and mortality associated with breast and ovarian cancer through the early detection of women at high risk. These women, once identified, can be targeted for more aggressive  
25 prevention programs. Screening is carried out by a variety of methods which include karyotyping, probe binding and DNA sequencing.

In DNA sequencing technology, genomic DNA is extracted from whole blood and the coding sequences of the BRCA1 gene are amplified. The coding sequences might be sequenced completely and the results are compared to the  
30 DNA sequence of the gene. Alternatively, the coding sequence of the sample gene may be compared to a panel of known mutations before completely sequencing the gene and comparing it to a normal sequence of the gene.

If a mutation in the BRCA1 coding sequence is found, it may be possible to provide the individual with increased expression of the gene through gene transfer therapy. It has been demonstrated that the gene transfer of the BRCA1 coding sequence into cancer cells inhibits their growth and reduces  
5 tumorigenesis of human cancer cells in nude mice. Jeffrey Holt and his colleagues conclude that the product of BRCA1 expression is a secreted tumor growth inhibitor, making BRCA1 an ideal gene for gene therapy studies. Transduction of only a moderate percentage of tumor cells apparently produces enough growth inhibitor to inhibit all tumor cells. Arteaga, CL, and JT Holt  
10 Cancer Research 56: 1098-1103 (1996), Holt, JT et al., Nature Genetics 12: 298-302 (1996).

The observation of Holt et al, that the BRCA1 growth inhibitor is a secreted protein leads to the possible use of injection of the growth inhibitor into the area of the tumor for tumor suppression.

15 The BRCA1 gene is divided into 24 separate exons. Exons 1 and 4 are noncoding, in that they are not part of the final functional BRCA1 protein product. The BRCA1 coding sequence spans roughly 5600 base pairs (bp). Each exon consists of 200-400 bp, except for exon 11 which contains about 3600 bp. To  
20 sequence the coding sequence of the BRCA1 gene, each exon is amplified separately and the resulting PCR products are sequenced in the forward and reverse directions. Because exon 11 is so large, we have divided it into twelve overlapping PCR fragments of roughly 350 bp each (segments "A" through "L" of BRCA1 exon 11).

Many mutations and polymorphisms have already been reported in the  
25 BRCA1 gene. A world wide web site has been built to facilitate the detection and characterization of alterations in breast cancer susceptibility genes. Such mutations in BRCA1 can be accessed through the Breast Cancer Information Core at: [http://www.nchgr.nih.gov/dir/lab\\_transfer/bic](http://www.nchgr.nih.gov/dir/lab_transfer/bic). This data site became publicly available on November 1, 1995. Friend, S. et al. Nature Genetics 11:238,  
30 (1995).

The genetics of Breast/Ovarian Cancer Syndrome is autosomal dominant with reduced penetrance. In simple terms, this means that the syndrome runs through families such that both sexes can be carriers (only women get the disease but men can pass it on), all generations will likely have breast/ovarian or both  
5 diseases and sometimes in the same individual, occasionally women carriers either die young before they have the time to manifest disease (and yet offspring get it) or they never develop breast or ovarian cancer and die of old age (the latter people are said to have "reduced penetrance" because they never develop cancer). Pedigree analysis and genetic counseling is absolutely essential to the proper  
10 workup of a family prior to any lab work.

Until now, only a single coding sequence for the BRCA1 gene has been available for comparison to patient samples. That sequence is available as GenBank Accession Number U14680. There is a need in the art, therefore, to have available a coding sequence which is the BRCA1 coding sequence found in  
15 the majority of the population, a "consensus coding sequence", BRCA1<sup>(omi1)</sup> Seq. ID. NO. 1. A consensus coding sequence will make it possible for true mutations to be easily identified or differentiated from polymorphisms. Identification of mutations of the BRCA1 gene and protein would allow more widespread diagnostic screening for hereditary breast and ovarian cancer than is currently  
20 possible. Two additional coding sequences have been isolated and characterize. The BRCA1<sup>(omi2)</sup> SEQ. ID. NO.: 3, and BRCA1<sup>(omi3)</sup> SEQ. ID. NO.:5 coding sequences also have utility in diagnosis, gene therapy and in making therapeutic BRCA1 protein.

A coding sequence of the BRCA1 gene which occurs most commonly in  
25 the human gene pool is provided. The most commonly occurring coding sequence more accurately reflects the most likely sequence to be found in a subject. Use of the coding sequence BRCA1<sup>(omi1)</sup> SEQ. ID. NO.: 1, rather than the previously published BRCA1 sequence, will reduce the likelihood of misinterpreting a "sequence variation" found in the population (i.e.  
30 polymorphism) with a pathologic "mutation" (i.e. causes disease in the individual or puts the individual at a high risk of developing the disease). With large interest in breast cancer predisposition testing, misinterpretation is

particularly worrisome. People who already have breast cancer are asking the clinical question: "is my disease caused by a heritable genetic mutation?" The relatives of the those with breast cancer are asking the question: "Am I also a carrier of the mutation my relative has? Thus, is my risk increased, and should I  
5 undergo a more aggressive surveillance program."

#### SUMMARY OF THE INVENTION

The present invention is based on the isolation of three coding sequences of the BRCA1 gene found in human individuals.

10 It is an object of the invention to provide the most commonly occurring coding sequence of the BRCA1 gene.

It is another object of this invention to provide two other coding sequences of BRCA1 gene.

15 It is another object of the invention to provide three protein sequences coded for by three of the coding sequences of the BRCA1 gene.

It is another object of the invention to provide a list of the codon pairs which occur at each of seven polymorphic points on the BRCA1 gene.

It is another object of the invention to provide the rates of occurrence for the codons.

20 It is another object of the invention to provide a method wherein BRCA1, or parts thereof, is amplified with one or more oligonucleotide primers.

It is another object of this invention to provide a method of identifying individuals who carry no mutation(s) of the BRCA1 coding sequence and therefore have no increased genetic susceptibility to breast or ovarian cancer  
25 based on their BRCA1 genes.

It is another object of this invention to provide a method of identifying a mutation leading to an increased genetic susceptibility to breast or ovarian cancer.

30 There is a need in the art for a sequence of the BRCA1 gene and for the protein sequence of BRCA1 as well as for an accurate list of codons which occur at polymorphic points on a sequence.

A person skilled in the art of genetic susceptibility testing will find the present invention useful for:

- 5 a) identifying individuals having a BRCA1 gene with no coding mutations, who therefore cannot be said to have an increased genetic susceptibility to breast or ovarian cancer from their BRCA1 genes;
- b) avoiding misinterpretation of polymorphisms found in the BRCA1 gene;
- 10 c) determining the presence of a previously unknown mutation in the BRCA1 gene.
- d) identifying a mutation which increases the genetic susceptibility to breast or ovarian cancer.
- e) probing a human sample of the BRCA1 gene.
- f) performing gene therapy.
- 15 g) for making a functioning tumor growth inhibitor protein coded for by one of the BRCA1<sup>omi</sup> genes.

#### BRIEF DESCRIPTION OF THE FIGURE

20 As shown in FIGURE 1, the alternative alleles at polymorphic (non-mutation causing variations) sites along a chromosome can be represented as a "haplotype" within a gene such as BRCA1. The BRCA1<sup>(omi1)</sup> haplotype is shown in Figure 1 with dark shading (encompassing the alternative alleles found at nucleotide sites 2201, 2430, 2731, 3232, 3667, 4427, and 4956). For comparison, the haplotype that is in GenBank is shown with no shading. As can be seen from the figure, the common "consensus" haplotype is found intact in five separate chromosomes labeled with the OMI symbol (numbers 1-5 from left to right). Two additional haplotypes (BRCA1<sup>(omi2)</sup>, and BRCA1<sup>(omi3)</sup>) are represented with mixed dark and light shading (numbers 7 and 9 from left to right). In total, 7 of 10 haplotypes along the BRCA1 gene are unique.

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## DETAILED DESCRIPTION OF THE INVENTION

### DEFINITIONS

The following definitions are provided for the purpose of understanding this  
5 invention.

"Breast and Ovarian cancer" is understood by those skilled in the art to include breast and ovarian cancer in women and also breast and prostate cancer in men. BRCA1 is associated genetic susceptibility to inherited breast and  
10 ovarian cancer in women and also breast and prostate cancer in men. Therefore, claims in this document which recite breast and/or ovarian cancer refer to breast, ovarian and prostate cancers in men and women.

"Coding sequence" or "DNA coding sequence" refers to those portions of a  
15 gene which, taken together, code for a peptide (protein), or which nucleic acid itself has function.

"Protein" or "peptide" refers to a sequence amino acids which has function.

20 "BRCA1<sup>(omi)</sup>" refers collectively to the "BRCA1<sup>(omi1)</sup>", "BRCA1<sup>(omi2)</sup>" and "BRCA1<sup>(omi3)</sup>" coding sequences.

"BRCA1<sup>(omi1)</sup>" refers to SEQ. ID. NO.: 1, a coding sequence for the BRCA1 gene. The coding sequence was found by end to end sequencing of BRCA1 alleles  
25 from individuals randomly drawn from a Caucasian population found to have no family history of breast or ovarian cancer. The sequenced gene was found not to contain any mutations. BRCA1<sup>(omi1)</sup> was determined to be a consensus sequence by calculating the frequency with which the coding sequence occurred among the sample alleles sequenced.

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"BRCA1<sup>(omi2)</sup>" and "BRCA1<sup>(omi3)</sup>" refer to SEQ. ID. NO.: 3, and SEQ. ID. NO.:  
5 respectively. They are two additional coding sequences for the BRCA1 gene

which were also isolated from individuals randomly drawn from a Caucasian population found to have no family history of breast or ovarian cancer. polymorphisms

5 "Primer" as used herein refers to a sequence comprising about 20 or more nucleotides of the BRCA1 gene.

"Genetic susceptibility" refers to the susceptibility to breast or ovarian cancer due to the presence of a mutation in the BRCA1 gene.

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A "target polynucleotide" refers to the nucleic acid sequence of interest *e.g.*, the BRCA1 encoding polynucleotide. Other primers which can be used for primer hybridization will be known or readily ascertainable to those of skill in the art.

15

"Consensus" means the most commonly occurring in the population.

"Consensus genomic sequence" means the allele of the target gene which occurs with the greatest frequency in a population of individuals having no family history of disease associated with the target gene.

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"Substantially complementary to" refers to a probe or primer sequences which hybridize to the sequences provided under stringent conditions and/or sequences having sufficient homology with BRCA1 sequences, such that the allele specific oligonucleotide probe or primers hybridize to the BRCA1 sequences to which they are complimentary.

25

"Haplotype" refers to a series of alleles within a gene on a chromosome.

30 "Isolated" as used herein refers to substantially free of other nucleic acids, proteins, lipids, carbohydrates or other materials with which they may be associated. Such association is typically either in cellular material or in a synthesis medium.

"Mutation" refers to a base change or a gain or loss of base pair(s) in a DNA sequence, which results in a DNA sequence which codes for a non-functioning protein or a protein with substantially reduced or altered function.

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"Polymorphism" refers to a base change which is not associated with known pathology.

"Tumor growth inhibitor protein" refers to the protein coded for by the BRCA1 gene. The functional protein is thought to suppress breast and ovarian tumor growth.

The invention in several of its embodiments includes:

1. An isolated consensus DNA sequence of the BRCA1 coding sequence as set forth in SEQ. ID. NO.: 1.
2. A consensus protein sequence of the BRCA1 protein as set forth in SEQ. ID. NO.: 2.
3. An isolated coding sequence of the BRCA1 gene as set forth in SEQ. ID. NO.: 3.
4. A protein sequence of the BRCA1 protein as set forth in SEQ. ID. NO.: 4.
5. An isolated coding sequence of the BRCA1 gene as set forth in SEQ. ID. NO.: 5.
6. A protein sequence of the BRCA1 protein as set forth in SEQ. ID. NO.: 6.
7. A BRCA1 gene with a BRCA1 coding sequence not associated with breast or ovarian cancer which comprises an alternative pair of codons, AGC



and AGT, which occur at position 2201 at frequencies of about 35-45%, and from about 55-65%, respectively.

5 8. A BRCA1 gene according to Claim 7 wherein AGC occurs at a frequency of about 40%.

9. A set of at least two alternative codon pairs which occur at polymorphic positions in a BRCA1 gene with a BRCA1 coding sequence not associated with breast or ovarian cancer, wherein codon pairs are selected  
10 from the group consisting of:

- AGC and AGT at position 2201;
- TTG and CTG at position 2430;
- CCG and CTG at position 2731;
- GAA and GGA at position 3232;
- 15 • AAA and AGA at position 3667;
- TCT and TCC at position 4427; and
- AGT and GGT at position 4956.

10. A set of at least two alternative codon pairs according to claim 9,  
20 wherein the codon pairs occur in the following frequencies, respectively, in a population of individuals free of disease:

- at position 2201, AGC and AGT occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 2430, TTG and CTG occur at frequencies from about  
25 35-45%, and from about 55-65%, respectively;
- at position 2731, CCG and CTG occur at frequencies from about 25-35%, and from about 65-75%, respectively;
- at position 3232, GAA and GGA occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- 30 • at position 3667, AAA and AGA occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 4427, TCT and TCC occur at frequencies from about 45-55%, and from about 45-55%, respectively; and
- at position 4956, AGT and GGT occur at frequencies from about

35-45%, and from about 55-65%, respectively.

11. A set according to Claim 10 which is at least three codon pairs.
- 5 12. A set according to Claim 10 which is at least four codon pairs.
13. A set according to Claim 10 which is at least five codon pairs.
14. A set according to Claim 10 which is at least six codon pairs.
- 10 15. A set according to Claim 10 which is at least seven codon pairs.
16. A method of identifying individuals having a BRCA1 gene with a BRCA1 coding sequence not associated with disease, comprising:
  - 15 (a) amplifying a DNA fragment of an individual's BRCA1 coding sequence using an oligonucleotide primer which specifically hybridizes to sequences within the gene;
  - (b) sequencing said amplified DNA fragment by dideoxy sequencing;
  - 20 (c) repeating steps (a) and (b) until said individual's BRCA1 coding sequence is completely sequenced;
  - (d) comparing the sequence of said amplified DNA fragment to a BRCA1<sup>(oml)</sup> DNA sequence, SEQ. ID. NO1, SEQ. ID. NO3, or SEQ. ID. NO5;
  - 25 (e) determining the presence or absence of each of the following polymorphic variation in said individual's BRCA1 coding sequence:
    - AGC and AGT at position 2201.
    - TTG and CTG at position 2430,
    - CCG and CTG at position 2731,
    - GAA and GGA at position 3232,
    - AAA and AGA at position 3667,
    - TCT and TCC at position 4427, and
    - AGT and GGT at position 4956;
- 30

- (f) determining any sequence differences between said individual's BRCA1 coding sequences and SEQ. ID. NO1, SEQ. ID. NO3, or SEQ. ID. NO5 wherein the presence of said polymorphic variations and the absence of a variation outside of positions 2201, 2430, 2731, 3232, 3667, 4427, and 4956, is correlated with an absence of increased genetic susceptibility to breast or ovarian cancer resulting from a BRCA1 mutation in the BRCA1 coding sequence.

17. A method of claim 16 wherein, codon variations occur at the following frequencies, respectively, in a population of individuals free of disease:

- at position 2201, AGC and AGT occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 2430, TTG and CTG occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 2731, CCG and CTG occur at frequencies from about 25-35%, and from about 65-75%, respectively;
- at position 3232, GAA and GGA occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 3667, AAA and AGA occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 4427, TCT and TCC occur at frequencies from about 45-55%, and from about 45-55%, respectively; and
- at position 4956, AGT and GGT occur at frequencies from about 35-45%, and from about 55-65%, respectively.

18. A method according to claim 16 wherein said oligonucleotide primer is labeled with a radiolabel, a fluorescent label a bioluminescent label, a chemiluminescent label, or an enzyme label.

19. A method of detecting a increased genetic susceptibility to breast and

ovarian cancer in an individual resulting from the presence of a mutation in the BRCA1 coding sequence, comprising:

- (a) amplifying a DNA fragment of an individual's BRCA1 coding sequence using an oligonucleotide primer which specifically hybridizes to sequences within the gene;
- (b) sequencing said amplified DNA fragment by dideoxy sequencing;
- (c) repeating steps (a) and (b) until said individual's BRCA1 coding sequence is completely sequenced;
- (d) comparing the sequence of said amplified DNA fragment to a BRCA1<sup>(oml)</sup> DNA sequence, SEQ. ID. NO1, SEQ. ID. NO3, or SEQ. ID. NO5;
- (e) determining any sequence differences between said individual's BRCA1 coding sequences and SEQ. ID. NO1, SEQ. ID. NO3, or SEQ. ID. NO5; to determine the presence or absence of base changes in said individual's BRCA1 coding sequence wherein a base change which is not any one of the following:
  - AGC and AGT at position 2201,
  - TTG and CTG at position 2430,
  - CCG and CTG at position 2731,
  - GAA and GGA at position 3232,
  - AAA and AGA at position 3667,
  - TCT and TCC at position 4427, and
  - AGT and GGT at position 4956 is correlated with the potential of increased genetic susceptibility to breast or ovarian cancer resulting from a BRCA1 mutation in the BRCA1 coding sequence.

20. A method of claim 19 wherein, codon variations occur at the following frequencies, respectively, in a population free of disease:

- at position 2201, AGC and AGT occur at frequencies from about 40%, and from about 55-65%, respectively;
- at position 2430, TTG and CTG occur at frequencies from about

35-45%, and from about 55-65%, respectively;

- at position 2731, CCG and CTG occur at frequencies from about 25-35%, and from about 65-75%, respectively;
- at position 3232, GAA and GGA occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 3667, AAA and AGA occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 4427, TCT and TCC occur at frequencies from about 45-55%, and from about 45-55%, respectively; and
- at position 4956, AGT and GGT occur at frequencies from about 35-45%, and from about 55-65%, respectively.

21. A method according to claim 19 wherein said oligonucleotide primer is labeled with a radiolabel, a fluorescent label a bioluminescent label, a chemiluminescent label, or an enzyme label.

22. A set of codon pairs, which occur at polymorphic positions in a BRCA1 gene with a BRCA1 coding sequence according to Claim 1, wherein said set of codon pairs is:

- AGC and AGT at position 2201;
- TTG and CTG at position 2430;
- CCG and CTG at position 2731;
- GAA and GGA at position 3232;
- AAA and AGA at position 3667;
- TCT and TCC at position 4427; and
- AGT and GGT at position 4956.

23. A set of at least two alternative codon pairs according to claim 22 wherein set of at least two alternative codon pairs occur at the following frequencies:

- at position 2201, AGC and AGT occur at frequencies of about 40%, and from about 55-65%, respectively;
- at position 2430, TTG and CTG occur at frequencies from about 35-45%, and from about 55-65%, respectively;

- at position 2731, CCG and CTG occur at frequencies from about 25-35%, and from about 65-75%, respectively;
- at position 3232, GAA and GGA occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- 5     • at position 3667, AAA and AGA occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 4427, TCT and TCC occur at frequencies from about 45-55%, and from about 45-55%, respectively; and
- 10    • at position 4956, AGT and GGT occur at frequencies from about 35-45%, and from about 55-65%, respectively.

24. A BRCA1 coding sequence according to claim 1 wherein the codon pairs occur at the following frequencies:

- 15     • at position 2201, AGC and AGT occur at frequencies of about 40%, and from about 55-65%, respectively;
- at position 2430, TTG and CTG occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 2731, CCG and CTG occur at frequencies from about 25-35%, and from about 65-75%, respectively;
- 20     • at position 3232, GAA and GGA occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 3667, AAA and AGA occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 4427, TCT and TCC occur at frequencies from about 45-55%, and from about 45-55%, respectively; and
- 25     • at position 4956, AGT and GGT occur at frequencies from about 35-45%, and from about 55-65%, respectively.

25. A method of determining the consensus genomic sequence or consensus coding sequence for a target gene, comprising:

- a) screening a number of individuals in a population for a family history which indicates inheritance of normal alleles for a target gene;
- b) isolating at least one allele of the target gene from individuals found to

have a family history which indicates inheritance of normal alleles for a target gene;

- c) sequencing each allele;
- d) comparing the nucleic acid sequence of the genomic sequence or of the coding sequence of each allele of the target gene to determine similarities and differences in the nucleic acid sequence; and
- e) determining which allele of the target gene occurs with the greatest frequency.

26. A method of performing gene therapy, comprising:

- a) transfecting cancer cell *in vivo* with an effective amount of a vector transformed with a BRCA1 coding sequences of SEQ. ID. NO.: 1, SEQ. ID. NO.: 3, or SEQ. ID. NO.: 5;
- b) allowing the cells to take up the vector, and
- c) measuring a reduction in tumor growth.

27. A method of performing protein therapy, comprising:

- a) injecting into a patient, an effective amount of BRCA1 tumor growth inhibiting protein of SEQ. ID. NO.: 2, SEQ. ID. NO.: 4, or SEQ. ID. NO.: 6;
- b) allowing the cells to take up the protein, and
- c) measuring a reduction in tumor growth.

## SEQUENCING

Any nucleic acid specimen, in purified or non-purified form, can be utilized as the starting nucleic acid or acids, providing it contains, or is suspected of containing, the specific nucleic acid sequence containing a polymorphic locus. Thus, the process may amplify, for example, DNA or RNA, including messenger RNA, wherein DNA or RNA may be single stranded or double stranded. In the event that RNA is to be used as a template, enzymes, and/or conditions optimal for reverse transcribing the template to DNA would be utilized. In addition, a DNA-RNA hybrid which contains one strand of each may be utilized. A mixture

of nucleic acids may also be employed, or the nucleic acids produced in a previous amplification reaction herein, using the same or different primers may be so utilized. See TABLE II. The specific nucleic acid sequence to be amplified, *i.e.*, the polymorphic locus, may be a fraction of a larger molecule or can be present initially as a discrete molecule, so that the specific sequence constitutes the entire nucleic acid. It is not necessary that the sequence to be amplified be present initially in a pure form; it may be a minor fraction of a complex mixture, such as contained in whole human DNA.

10 DNA utilized herein may be extracted from a body sample, such as blood, tissue material and the like by a variety of techniques such as that described by Maniatis, *et. al.* in *Molecular Cloning: A Laboratory Manual*, Cold Spring Harbor, NY, p 280-281, 1982). If the extracted sample is impure, it may be treated before amplification with an amount of a reagent effective to open the cells, or animal  
15 cell membranes of the sample, and to expose and/or separate the strand(s) of the nucleic acid(s). This lysing and nucleic acid denaturing step to expose and separate the strands will allow amplification to occur much more readily.

The deoxyribonucleotide triphosphates dATP, dCTP, dGTP, and dTTP are added to the synthesis mixture, either separately or together with the primers, in  
20 adequate amounts and the resulting solution is heated to about 90°-100°C from about 1 to 10 minutes, preferably from 1 to 4 minutes. After this heating period, the solution is allowed to cool, which is preferable for the primer hybridization. To the cooled mixture is added an appropriate agent for effecting the primer extension reaction (called herein "agent for polymerization"), and the reaction is  
25 allowed to occur under conditions known in the art. The agent for polymerization may also be added together with the other reagents if it is heat stable. This synthesis (or amplification) reaction may occur at room temperature up to a temperature above which the agent for polymerization no longer functions. Thus, for example, if DNA polymerase is used as the agent, the  
30 temperature is generally no greater than about 40°C. Most conveniently the reaction occurs at room temperature.

The primers used to carry out this invention embrace oligonucleotides of



sufficient length and appropriate sequence to provide initiation of polymerization. Environmental conditions conducive to synthesis include the presence of nucleoside triphosphates and an agent for polymerization, such as DNA polymerase, and a suitable temperature and pH. The primer is preferably  
5 single stranded for maximum efficiency in amplification, but may be double stranded. If double stranded, the primer is first treated to separate its strands before being used to prepare extension products. The primer must be sufficiently long to prime the synthesis of extension products in the presence of the inducing agent for polymerization. The exact length of primer will depend on many  
10 factors, including temperature, buffer, and nucleotide composition. The oligonucleotide primer typically contains 12-20 or more nucleotides, although it may contain fewer nucleotides.

Primers used to carry out this invention are designed to be substantially complementary to each strand of the genomic locus to be amplified. This means  
15 that the primers must be sufficiently complementary to hybridize with their respective strands under conditions which allow the agent for polymerization to perform. In other words, the primers should have sufficient complementarity with the 5' and 3' sequences flanking the mutation to hybridize therewith and permit amplification of the genomic locus.

20 Oligonucleotide primers of the invention are employed in the amplification process which is an enzymatic chain reaction that produces exponential quantities of polymorphic locus relative to the number of reaction steps involved. Typically, one primer is complementary to the negative (-) strand of the polymorphic locus and the other is complementary to the positive  
25 (+) strand. Annealing the primers to denatured nucleic acid followed by extension with an enzyme, such as the large fragment of DNA polymerase I (Klenow) and nucleotides, results in newly synthesized + and - strands containing the target polymorphic locus sequence. Because these newly synthesized sequences are also templates, repeated cycles of denaturing, primer  
30 annealing, and extension results in exponential production of the region (*i.e.*, the target polymorphic locus sequence) defined by the primers. The product of the chain reaction is a discrete nucleic acid duplex with termini corresponding to the

ends of the specific primers employed.

The oligonucleotide primers of the invention may be prepared using any suitable method, such as conventional phosphotriester and phosphodiester methods or automated embodiments thereof. In one such automated  
5 embodiment, diethylphosphoramidites are used as starting materials and may be synthesized as described by Beaucage, et al., Tetrahedron Letters, 22:1859-1862, 1981. One method for synthesizing oligonucleotides on a modified solid support is described in U.S. Patent No. 4,458,066.

10 The agent for polymerization may be any compound or system which will function to accomplish the synthesis of primer extension products, including enzymes. Suitable enzymes for this purpose include, for example, *E. coli* DNA polymerase I, Klenow fragment of *E. coli* DNA polymerase, polymerase muteins,  
15 reverse transcriptase, other enzymes, including heat-stable enzymes (*e.i.*, those enzymes which perform primer extension after being subjected to temperatures sufficiently elevated to cause denaturation), such as *Taq* polymerase. Suitable enzyme will facilitate combination of the nucleotides in the proper manner to form the primer extension products which are complementary to each polymorphic locus nucleic acid strand. Generally, the synthesis will be initiated  
20 at the 3' end of each primer and proceed in the 5' direction along the template strand, until synthesis terminates, producing molecules of different lengths.

The newly synthesized strand and its complementary nucleic acid strand will form a double-stranded molecule under hybridizing conditions described above and this hybrid is used in subsequent steps of the process. In the next step,  
25 the newly synthesized double-stranded molecule is subjected to denaturing conditions using any of the procedures described above to provide single-stranded molecules.

The steps of denaturing, annealing, and extension product synthesis can be repeated as often as needed to amplify the target polymorphic locus nucleic acid  
30 sequence to the extent necessary for detection. The amount of the specific nucleic acid sequence produced will accumulate in an exponential fashion. Amplification is described in PCR, A Practical Approach, ILR Press, Eds. M. J.

McPherson, P. Quirke, and G. R. Taylor, 1992.

The amplification products may be detected by Southern blots analysis, without using radioactive probes. In such a process, for example, a small sample of DNA containing a very low level of the nucleic acid sequence of the polymorphic locus is amplified, and analyzed via a Southern blotting technique or similarly, using dot blot analysis. The use of non-radioactive probes or labels is facilitated by the high level of the amplified signal. Alternatively, probes used to detect the amplified products can be directly or indirectly detectably labeled, for example, with a radioisotope, a fluorescent compound, a bioluminescent compound, a chemiluminescent compound, a metal chelator or an enzyme. Those of ordinary skill in the art will know of other suitable labels for binding to the probe, or will be able to ascertain such, using routine experimentation.

Sequences amplified by the methods of the invention can be further evaluated, detected, cloned, sequenced, and the like, either in solution or after binding to a solid support, by any method usually applied to the detection of a specific DNA sequence such as PCR, oligomer restriction, (Saiki, *et.al.*, *Bio/Technology*, 3:1008-1012, 1985), allele-specific oligonucleotide (ASO) probe analysis (Conner, *et. al.*, *Proc. Natl. Acad. Sci. U.S.A.*, 80:278, 1983), oligonucleotide ligation assays (OLAs) (Landgren, *et. al.*, *Science*, 241:1007, 1988), and the like. Molecular techniques for DNA analysis have been reviewed (Landgren, *et. al.*, *Science*, 242:229-237, 1988).

Preferably, the method of amplifying is by PCR, as described herein and as is commonly used by those of ordinary skill in the art. Alternative methods of amplification have been described and can also be employed as long as the BRCA1 locus amplified by PCR using primers of the invention is similarly amplified by the alternative means. Such alternative amplification systems include but are not limited to self-sustained sequence replication, which begins with a short sequence of RNA of interest and a T7 promoter. Reverse transcriptase copies the RNA into cDNA and degrades the RNA, followed by reverse transcriptase polymerizing a second strand of DNA. Another nucleic acid amplification technique is nucleic acid sequence-based amplification (NASBA) which uses reverse transcription and T7 RNA polymerase and

incorporates two primers to target its cycling scheme. NASBA can begin with either DNA or RNA and finish with either, and amplifies to  $10^8$  copies within 60 to 90 minutes. Alternatively, nucleic acid can be amplified by ligation activated transcription (LAT). LAT works from a single-stranded template with a single  
5 primer that is partially single-stranded and partially double-stranded. Amplification is initiated by ligating a cDNA to the promoter oligonucleotide and within a few hours, amplification is  $10^8$  to  $10^9$  fold. Another amplification system useful in the method of the invention is the QB Replicase System. The QB replicase system can be utilized by attaching an RNA sequence called MDV-1  
10 to RNA complementary to a DNA sequence of interest. Upon mixing with a sample, the hybrid RNA finds its complement among the specimen's mRNAs and binds, activating the replicase to copy the tag-along sequence of interest. Another nucleic acid amplification technique, ligase chain reaction (LCR), works by using two differently labeled halves of a sequence of interest which are  
15 covalently bonded by ligase in the presence of the contiguous sequence in a sample, forming a new target. The repair chain reaction (RCR) nucleic acid amplification technique uses two complementary and target-specific oligonucleotide probe pairs, thermostable polymerase and ligase, and DNA nucleotides to geometrically amplify targeted sequences. A 2-base gap separates  
20 the oligonucleotide probe pairs, and the RCR fills and joins the gap, mimicking DNA repair. Nucleic acid amplification by strand displacement activation (SDA) utilizes a short primer containing a recognition site for *hincII* with short overhang on the 5' end which binds to target DNA. A DNA polymerase fills in the part of the primer opposite the overhang with sulfur-containing adenine  
25 analogs. *HincII* is added but only cuts the unmodified DNA strand. A DNA polymerase that lacks 5' exonuclease activity enters at the site of the nick and begins to polymerize, displacing the initial primer strand downstream and building a new one which serves as more primer. SDA produces greater than  $10^7$ -fold amplification in 2 hours at 37°C. Unlike PCR and LCR, SDA does not  
30 require instrumented Temperature cycling.

Another method is a process for amplifying nucleic acid sequences from a DNA or RNA template which may be purified or may exist in a mixture of

nucleic acids. The resulting nucleic acid sequences may be exact copies of the template, or may be modified. The process has advantages over PCR in that it increases the fidelity of copying a specific nucleic acid sequence, and it allows one to more efficiently detect a particular point mutation in a single assay. A target  
5 nucleic acid is amplified enzymatically while avoiding strand displacement. Three primers are used. A first primer is complementary to the first end of the target. A second primer is complementary to the second end of the target. A third primer which is similar to the first end of the target and which is substantially complementary to at least a portion of the first primer such that  
10 when the third primer is hybridized to the first primer, the position of the third primer complementary to the base at the 5' end of the first primer contains a modification which substantially avoids strand displacement. This method is detailed in U.S. Patent 5,593,840 to Bhatnagar et al. 1997. Although PCR is the preferred method of amplification if the invention, these other methods can also  
15 be used to amplify the BRCA1 locus as described in the method of the invention.

The BRCA1<sup>(omi)</sup> DNA coding sequences were obtained by end to end sequencing of the BRCA1 alleles of five subjects in the manner described above followed by analysis of the data obtained. The data obtained provided us with the opportunity to evaluate seven previously published polymorphisms and to  
20 affirm or correct where necessary, the frequency of occurrence of alternative codons.

#### GENE THERAPY

The coding sequences can be used for gene therapy.

A variety of methods are known for gene transfer, any of which might be  
25 available for use.

##### *Direct injection of Recombinant DNA in vivo*

1. Direct injection of "naked" DNA directly with a syringe and needle into a specific tissue, infused through a vascular bed, or transferred through a catheter into endothelial cells.
- 30 2. Direct injection of DNA that is contained in artificially generated lipid vesicles.
3. Direct injection of DNA conjugated to a targeting structure, such as an

antibody.

4. Direct injection by particle bombardment, where the DNA is coated onto gold particles and shot into the cells.

5 *Human Artificial Chromosomes*

This novel gene delivery approach involves the use of human chromosomes that have been striped down to contain only the essential components for replication and the genes desired for transfer.

10 *Receptor-Mediated Gene Transfer*

DNA is linked to a targeting molecule that will bind to specific cell-surface receptors, inducing endocytosis and transfer of the DNA into mammalian cells. One such technique uses poly-L-lysine to link asialoglycoprotein to DNA. An adenovirus is also added to the complex to disrupt the lysosomes and thus allow the DNA to avoid degradation and move to the nucleus. Infusion of these particles intravenously has resulted in gene transfer into hepatocytes.

RECOMBINANT VIRUS VECTORS

Several vectors are used in gene therapy. Among them are the Moloney Murine Leukemia Virus (MoMLV) Vectors, the adenovirus vectors, the adeno-Associated Virus (AAV) vectors, the herpes simplex virus (HSV) vectors, the poxvirus vectors, and human immunodeficiency virus (HIV) vectors,

GENE REPLACEMENT AND REPAIR

25 The ideal genetic manipulation for treatment of a genetic disease would be the actual replacement of the defective gene with a normal copy of the gene. Homologous recombination is the term used for switching out a section of DNA and replacing it with a new piece. By this technique, the defective gene can be replaced with a normal gene which expresses a functioning BRCA1 tumor growth inhibitor protein.

30 A complete description of gene therapy can also be found in "Gene Therapy A Primer For Physicians 2d Ed. by Kenneth W. Culver, M.D. Publ. Mary Ann

Liebert Inc. (1996). Two Gene Therapy Protocols for BRCA1 are approved by the Recombinant DNA Advisory Committee for Jeffrey T. Holt *et al.*. They are listed as 9602-148, and 9603-149 and are available from the NIH. The isolated BRCA1 gene can be synthesized or constructed from amplification products and inserted  
5 into a vector such as the LXS vector.

The BRCA1 amino acid and nucleic acid sequence may be used to make diagnostic probes and antibodies. Labeled diagnostic probes may be used by any hybridization method to determine the level of BRCA1 protein in serum or lysed cell suspension of a patient, or solid surface cell sample.

10 The BRCA1 amino acid sequence may be used to provide a level of protection for patients against risk of breast or ovarian cancer or to reduce the size of a tumor. Methods of making and extracting proteins are well known. Itakura *et al.* U.S. Patents 4,704,362, 5, 221, 619, and 5,583,013. BRCA1 has been shown to be secreted. Jensen, R.A. *et al. Nature Genetics* 12: 303-308 (1996).

15

#### EXAMPLE 1

#### Determination Of The Coding Sequence Of A BRCA1<sup>(omi)</sup> Gene From Five Individuals

#### 20 MATERIALS AND METHODS

Approximately 150 volunteers were screened in order to identify individuals with no cancer history in their immediate family (i.e. first and second degree relatives). Each person was asked to fill out a hereditary cancer prescreening questionnaire See TABLE I below. Five of these were randomly  
25 chosen for end-to-end sequencing of their BRCA1 gene. A first degree relative is a parent, sibling, or offspring. A second degree relative is an aunt, uncle, grandparent, grandchild, niece, nephew, or half-sibling.

TABLE I

## Hereditary Cancer Pre-Screening Questionnaire

Part A: Answer the following questions about your family

- 5 1. To your knowledge, has anyone in your family been diagnosed with a very specific hereditary colon disease called Familial Adenomatous Polyposis (FAP)?
2. To your knowledge, have you or any aunt had breast cancer diagnosed before the age 35?
3. Have you had Inflammatory Bowel Disease, also called Crohn's Disease or Ulcerative Colitis, for more than 7 years?

Part B: Refer to the list of cancers below for your responses only to questions in Part B

- 10
- |                    |                    |                   |
|--------------------|--------------------|-------------------|
| Bladder Cancer     | Lung Cancer        | Pancreatic Cancer |
| Breast Cancer      | Gastric Cancer     | Prostate Cancer   |
| Colon Cancer       | Malignant Melanoma | Renal Cancer      |
| Endometrial Cancer | Ovarian Cancer     | Thyroid Cancer    |
- 15 4. Have your mother or father, your sisters or brothers or your children had any of the listed cancers?
- 5 Have there been diagnosed in your mother's brothers or sisters, or your mother's parents more than one of the cancers in the above list?
- 6 Have there been diagnosed in your father's brothers or sisters, or your father's parents more than one of the cancers in the above list?

Part C: Refer to the list of relatives below for responses only to questions in Part C

- 20
- |                          |   |
|--------------------------|---|
| You                      | Your mother   |
| Your sisters or brothers | Your mother's sisters or brothers (maternal aunts and uncles) |
| Your children            | Your mother's parents (maternal grandparents)                 |
- 7 Have there been diagnosed in these relatives 2 or more identical types of cancer?
- 25 Do not count "simple" skin cancer, also called basal cell or squamous cell skin cancer.
- 8 Is there a total of 4 or more of any cancers in the list of relatives above other than "simple" skin cancers?

Part D: Refer to the list of relatives below for responses only to questions in Part D.

- 30
- |                          |   |
|--------------------------|---|
| You                      | Your father   |
| Your sisters or brothers | Your father's sisters or brothers (paternal aunts and uncles) |
| Your children            | Your father's parents (paternal grandparents)                 |
9. Have there been diagnosed in these relatives 2 or more identical types of cancer?
- Do not count "simple" skin cancer, also called basal cell or squamous cell skin cancer.
10. Is there a total of 4 or more of any cancers in the list of relatives above other than "simple" skin cancers?
- 35



Genomic DNA was isolated from white blood cells of five subjects selected from analysis of their answers to the questions above. Dideoxy sequence analysis was performed following polymerase chain reaction amplification.

5 All exons of the BRCA1 gene were subjected to direct dideoxy sequence analysis by asymmetric amplification using the polymerase chain reaction (PCR) to generate a single stranded product amplified from this DNA sample. Shuldiner, *et al.*, Handbook of Techniques in Endocrine Research, p. 457-486, DePablo, F., Scanes, C., eds., Academic Press, Inc., 1993. Fluorescent dye was attached for automated sequencing using the Taq Dye Terminator® Kit (Perkin-  
10 Elmer cat# 401628). DNA sequencing was performed in both forward and reverse directions on an Applied Biosystems, Inc. (ABI) automated Model 377® sequencer. The software used for analysis of the resulting data was Sequence Navigator® software purchased through ABI.

15 1. Polymerase Chain Reaction (PCR) Amplification

Genomic DNA (100 nanograms) extracted from white blood cells of five subjects. Each of the five samples was sequenced end to end. Each sample was amplified in a final volume of 25 microliters containing 1 microliter (100 nanograms) genomic DNA, 2.5 microliters 10X PCR buffer (100 mM Tris, pH 8.3,  
20 500 mM KCl, 1.2 mM MgCl<sub>2</sub>), 2.5 microliters 10X dNTP mix (2 mM each nucleotide), 2.5 microliters forward primer, 2.5 microliters reverse primer, and 1 microliter Taq polymerase (5 units), and 13 microliters of water.

The primers in Table II, below were used to carry out amplification of the various sections of the BRCA1 gene samples. The primers were synthesized on  
25 an DNA/RNA Model 394® Synthesizer.

TABLE II  
BRCA1 PRIMERS AND SEQUENCING DATA

	EXON	SEQUENCE	SEQ.ID NO.	MER	Mg <sup>++</sup>	SIZE
5	EXON2	2F 5' GAA GTT GTC ATT TTA TAA ACC TTT-3' 2R 5' TGT CTT TTC TTC OCT AGT ATG T-3'	7 8	24 22	1.6	~275
10	EXON3	3F 5' TCC TGA CAC AGC AGA CAT TTA-3' 3R 5' TTG GAT TTT CGT TCT CAC TTA-3'	9 10	21 21	1.4	~375
	EXON5	5F 5' CTC TTA AGG GCA GTT GTG AG-3' 5R 5' TTC CTA CTG TGG TTG CTT CC	11 12	20 20 <sup>1</sup>	1.2	~275
15	EXON6	6/7F 5' CTT ATT TTA GTG TCC TTA AAA GG-3' 6R 5' TTT CAT GGA CAG CAC TTG AGT G-3'	13 14	23 22	1.6	~250
	EXON7	7F 5' CAC AAC AAA GAG CAT ACA TAG GG-3' 6/7R 5' TCG GGT TCA CTC TGT AGA AG-3'	15 16	23 20	1.6	~275
20	EXON8	8F1 5' TTC TCT TCA GGA GGA AAA GCA-3' 8R1 5' GCT GGC TAC CAC AAA TAC AAA-3'	17 18	21 21	1.2	~270
25	EXON9	9F 5' CCA CAG TAG ATG CTC AGT AAATA-3' 9R 5' TAG GAA AAT ACC AGC TTC ATA GA-3'	19 20	23 23	1.2	~250
	EXON10	10F 5' TGG TCA GCT TTC TGT AAT CG-3' 10R 5' GTA TCT ACC CAC TCT CTT CTT CAG-3'	21 22	20 24	1.6	~250
30	EXON11A11AF	5' CCA CCT CCA AGG TGT ATC A-3' 11AR 5' TGT TAT GTT GGC TCC TTG CT-3'	23 24	19 20	1.2	372
	EXON11B11BF1	5' CAC TAA AGA CAG AAT GAA TCT A-3; 11BR1 5' GAA GAA CCA GAA TAT TCA TCT A-3'	25 26	21 21	1.2	~400
35	EXON11C11CF1	5' TGA TGG GGA GTC TGA ATC AA-3' 11CR1 5' TCT GCT TTC TTG ATA AAA TCC T-3'	27 28	20 22	1.2	~400
40	EXON11D11DF1	5' AGC GTC CCC TCA CAA ATA AA-3' 11DR1 5' TCA AGC GCA TGA ATA TGC CT-3'	29 30	20 20	1.2	~400
	EXON11E11EF	5' GTA TAA GCA ATA TGG AAC TCG A-3' 11ER 5' TTA AGT TCA CTG GTA TTT GAA CA-3'	31 32	22 23	1.2	388
45	EXON11F11FF	5' GAC AGC GAT ACT TTC CCA GA-3' 11FR 5' TGG AAC AAC CAT GAA TTA GTC-3'	33 34	20 21	1.2	382
	EXON11G11GF	5' GGA AGT TAG CAC TCT AGG GA-3' 11GR 5' GCA GTG ATA TTA ACT GTC TGT A-3'	35 36	20 22	1.2	423

<sup>1</sup> M13 tailed

	EXON 11H11HF	5'	TGG	GTC	CTT	AAA	GAA	ACA	AAGT-3'	37	22	1.2	366
			11HR	5'	TCA	GGT	GAC	ATT	GAA TCT TCC-3'	38	21		
5	EXON 11I 11IF	5'	CCA	CTT	TTT	CCC	ATC	AAG	TCA-3'	39	21	1.2	377
			11IR	5'	TCA	GGA	TGC	TTA	CAA TTA CTT C-3'	40	21		
	EXON 11J11JF	5'	CAA	AAT	TGA	ATG	CTA	TGC	TTA GA-3'	41	23	1.2	377
			11JR	5'	TCG	GTA	ACC	CTG	AGC CAA AT-3'	42	20		
10	EXON 11K11KF	5'	GCA	AAAGCG	TCC	AGA	AAG	GA-3'		43	20	1.2	396
			11KR-15'	TAT	TTG	CAG	TCA	AGT	CTT CCA A-3'	44	22		
15	EXON 11L 11LF-1	5'	GTA	ATA	TTG	GCA	AAG	GCA	TCT-3'	45	22	1.2	360
			11LR	5'	TAA	AAT	GTG	CTC	CCC AAA AGC A-3'	46	22		
	EXON 12 12F	5'	GTC	CTG	CCA	ATG	AGA	AGA	AA-3'	47	20	1.2	-300
			12R	5'	TGT	CAG	CAA	ACC	TAA GAA TGT-3'	48	21		
20	EXON 13 13F	5'	AAT	GGA	AAG	CTT	CTC	AAAGTA-3'		49	21	1.2	-325
			13R	5'	ATG	TTG	GAG	CTA	GGT CCT TAC-3'	50	21		
	EXON 14 14F	5'	CTA	ACC	TGA	ATT	ATC	ACT	ATC A-3'	51	22	1.2	-310
			14R	5'	GTG	TAT	AAATGC	CTG	TAT GCA-3'	52	21		
25	EXON 15 15F	5'	TGG	CTG	CCC	AGG	AAG	TAT	G-3'	53	19	1.2	-375
			15R	5'	AAC	CAG	AAT	ATC	TTT ATG TAG GA-3'	54	23		
30	EXON 16 16F	5'	AAT	TCT	TAA	CAG	AGA	CCA	GAA C-3'	55	22	1.6	-550
			16R	5'	AAA	ACT	CTT	TCC	AGA ATG TTG T-3'	56	22		
	EXON 17 17F	5'	GTG	TAG	AAC	GTG	CAG	GAT	TG-3'	57	20	1.2	-275
			17R	5'	TCG	CCT	CAT	GTG	GTT TTA-3'	58	18		
35	EXON 18 18F	5'	GGC	TCT	TTA	GCT	TCT	TAG	GAC-3'	59	21	1.2	-350
			18R	5'	GAG	ACC	ATT	TTC	CCA GCA TC-3'	60	20		
	EXON 19 19F	5'	CTG	TCA	TTC	TTC	CTG	TGC	TC-3'	61	20	1.2	-250
			19R	5'	CAT	TGT	TAA	GGA	AAG TGG TGC-3'	62	21		
40	EXON 20 20F	5'	ATA	TGA	CGT	GTC	TGC	TCC	AC-3'	63	20	1.2	-425
			20R	5'	GGG	AAT	CCA	AAT	TAC ACA GC-3'	64	20		
45	EXON 21 21F	5'	AAG	CTC	TTC	CTT	TTT	GAA	AGT C-3'	65	22	1.6	-300
			21R	5'	GTA	GAG	AAA	TAG	AAT AGC CTC T-3'	66	22		
	EXON 22 22F	5'	TCC	CAT	TGA	GAG	GTC	TTG	CT-3'	67	20	1.6	-300
			22R	5'	GAG	AAG	ACT	TCT	GAG GCT AC-3'	68	20		
50	EXON 23 23F-1	5'	TGA	AGT	GAC	AGT	TCC	AGT	AGT-3'	69	21	1.2	-250
			23R-1	5'	CAT	TTT	AGC	CAT	TCA TTC AAC AA-3'	70	23		
55	EXON 24 24F	5'	ATG	AAT	TGA	CAC	TAA	TCT	CTG C-3'	71	22	1.4	-285
			24R	5'	GTA	GCC	AGG	ACA	GTA GAA GGA-3'	72	21		

Thirty-five cycles were performed, each consisting of denaturing (95°C; 30 seconds), annealing (55°C; 1 minute), and extension (72°C; 90 seconds), except during the first cycle in which the denaturing time was increased to 5 minutes, and during the last cycle in which the extension time was increased to 5 minutes.

- 5        PCR products were purified using Qia-quick® PCR purification kits (Qiagen cat# 28104; Chatsworth, CA). Yield and purity of the PCR product determined spectrophotometrically at OD<sub>260</sub> on a Beckman DU 650 spectrophotometer.

## 2.        Dideoxy Sequence Analysis

- Fluorescent dye was attached to PCR products for automated sequencing using the Taq  
10    Dye Terminator® Kit (Perkin-Elmer cat# 401628). DNA sequencing was performed in both forward and reverse directions on an Applied Biosystems, Inc. (ABI) Foster City, CA., automated Model 377® sequencer. The software used for analysis of the resulting data was "Sequence Navigator® software" purchased through ABI.

## 3.        RESULTS

- 15    Differences in the nucleic acids of the ten alleles from five individuals were found in seven locations on the gene. The changes and their positions are found on TABLE III, below.

TABLE III  
PANEL TYPING

5	AMINO ACID CHANGE	NUCLEOTIDE CHANGE						FREQUENCY
			1	2	3	4	5	
10	SER(SER) (694)	11E	C/C	C/T	C/T	T/T	T/T	0.4 C 0.6 T
	LEU(LEU) (771)	11F	T/T	C/T	C/T	C/C	C/C	0.4 T 0.6 C
15	PRO(LEU) (871)	11G	C/T	C/T	C/T	T/T	T/T	0.3 C 0.7 T
	GLU(GLY) (1038)	11I	A/A	A/G	A/G	G/G	G/G	0.4 A 0.6 G
20	LYS(ARG) (1183)	11J	A/A	A/G	A/G	G/G	G/G	0.4 A 0.6 G
	SER(SER) (1436)	13	T/T	T/T	T/C	C/C	C/C	0.5 T 0.5 C
25	SER(GLY) (1613)	16	A/A	A/G	A/G	G/G	G/G	0.4 A 0.6 G

Tables 3 and 4 depict one aspect of the invention, sets of at least two alternative codon pairs wherein the codon pairs occur in the following frequencies, respectively, in a population of individuals free of disease:

- at position 2201, AGC and AGT occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 2430, TTG and CTG occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 2731, CCG and CTG occur at frequencies from about 25-35%, and from about 65-75%, respectively;
- at position 3232, GAA and GGA occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 3667, AAA and AGA occur at frequencies from about 35-45%, and from about 55-65%, respectively;
- at position 4427, TCT and TCC occur at frequencies from about 45-55%, and from about 45-55%, respectively; and
- at position 4956, AGT and GGT occur at frequencies from about 35-45%, and from about 55-65%, respectively.

The data show that for each of the samples. The BRCA1 gene is identical except in the region of seven polymorphisms. These polymorphic regions, together with their locations, the amino acid groups of each codon, the frequency of their occurrence and the amino acid coded for by each codon are found in TABLE IV below.

TABLE IV  
CODON AND BASE CHANGES IN SEVEN POLYMORPHIC SITES OF BRCA1 GENE

SAMPLE NAME	BASE CHANGE	POSITION nt/aa	EXON	CODON CHANGE	AA CHANGE	PUBLISHED FREQUENCY <sup>2</sup>	FREQUENCY IN THIS STUDY
2,3,4,5	C-T	2201/694	11E	AGC(AGT)	SER-SER	UNPUBLISHED	C=40%
2,3,4,5	T-C	2430/771	11F	TTG(CTG)	LEU-LEU	T=67% <sup>13</sup>	T=40%
1,2,3,4,5	C-T	2731/871	11G	CCG(CTG)	PRO-LEU	C=34% <sup>12</sup>	C=30%



## EXAMPLE 2

Determination Of A Individual Using BRCA1<sup>(OMI)</sup> And The Seven Polymorphisms For Reference

5 A person skilled in the art of genetic susceptibility testing will find the present invention useful for:

- a) identifying individuals having a BRCA1 gene, who are therefore have no elevated genetic susceptibility to breast or ovarian cancer from a BRCA1 mutation;
  - b) avoiding misinterpretation of polymorphisms found in the
- 10 BRCA1 gene;

Sequencing is carried out as in EXAMPLE 1 using a blood sample from the patient in question. However, a BRCA1<sup>(omi)</sup> sequence is used for reference and the polymorphic sites are compared to the nucleic acid sequences listed above for codons at each polymorphic site. A sample is one which compares to a BRCA1<sup>(omi)</sup> sequence and

15 contains one of the base variations which occur at each of the polymorphic sites. The codons which occur at each of the polymorphic sites are paired here reference.

- AGC and AGT at position 2201,
- TTG and CTG at position 2430,
- CCG and CTG at position 2731,
- 20 • GAA and GGA at position 3232,
- AAA and AGA at position 3667,
- TCT and TCC at position 4427, and
- AGT and GGT at position 4956.

The availability of these polymorphic pairs provides added assurance that one skilled in

25 the art can correctly interpret the polymorphic variations without mistaking a variation for a mutation.

Exon 11 of the BRCA1 gene is subjected to direct dideoxy sequence analysis by asymmetric amplification using the polymerase chain reaction (PCR) to generate a single stranded product amplified from this DNA sample. Shuldiner, *et al.*, Handbook

30 of Techniques in Endocrine Research, p. 457-486, DePablo, F., Scanes, C., eds., Academic Press, Inc., 1993. Fluorescent dye is attached for automated sequencing using the Taq Dye Terminator® Kit (Perkin-Elmer cat# 401628). DNA sequencing is performed in



both forward and reverse directions on an Applied Biosystems, Inc. (ABI) automated Model 377<sup>®</sup> sequencer. The software used for analysis of the resulting data is "Sequence Navigator<sup>®</sup> software" purchased through ABI.

5 1. Polymerase Chain Reaction (PCR) Amplification

Genomic DNA (100 nanograms) extracted from white blood cells of the subject is amplified in a final volume of 25 microliters containing 1 microliter (100 nanograms) genomic DNA, 2.5 microliters 10X PCR buffer (100 mM Tris, pH 8.3, 500 mM KCl, 1.2 mM MgCl<sub>2</sub>), 2.5 microliters 10X dNTP mix (2 mM each nucleotide), 2.5 microliters  
10 forward primer (BRCA1-11K-F, 10 micromolar solution), 2.5 microliters reverse primer (BRCA1-11K-R, 10 micromolar solution), and 1 microliter Taq polymerase (5 units), and 13 microliters of water.

The PCR primers used to amplify a patient's sample BRCA1 gene are listed in Table II. The primers were synthesized on an DNA/RNA Model 394<sup>®</sup> Synthesizer. Thirty-five  
15 cycles are of amplification are performed, each consisting of denaturing (95°C; 30 seconds), annealing (55°C; 1 minute), and extension (72°C; 90 seconds), except during the first cycle in which the denaturing time is increased to 5 minutes, and during the last cycle in which the extension time is increased to 5 minutes.

PCR products are purified using Qia-quick<sup>®</sup> PCR purification kits (Qiagen, cat# 28104; Chatsworth, CA). Yield and purity of the PCR product determined  
20 spectrophotometrically at OD<sub>260</sub> on a Beckman DU 650 spectrophotometer.

2. Dideoxy Sequence Analysis

Fluorescent dye is attached to PCR products for automated sequencing using the Taq  
25 Dye Terminator<sup>®</sup> Kit (Perkin-Elmer cat# 401628). DNA sequencing is performed in both forward and reverse directions on an Applied Biosystems, Inc. (ABI) Foster City, CA., automated Model 377<sup>®</sup> sequencer. The software used for analysis of the resulting data is "Sequence Navigator<sup>®</sup> software" purchased through ABI. The BRCA1<sup>(omil)</sup> SEQ. ID. NO.:1 sequence is entered into the Sequence Navigator<sup>®</sup> software as the Standard for  
30 comparison. The Sequence Navigator<sup>®</sup> software compares the sample sequence to the BRCA1<sup>(omil)</sup> SEQ. ID. NO.:1 standard, base by base. The Sequence Navigator<sup>®</sup> software

highlights all differences between the BRCA1<sup>(omi1)</sup> SEQ. ID. NO.:1 DNA sequence and the patient's sample sequence.

A first technologist checks the computerized results by comparing visually the BRCA1<sup>(omi1)</sup> SEQ. ID. NO.:1 standard against the patient's sample, and again highlights any differences between the standard and the sample. The first primary technologist then interprets the sequence variations at each position along the sequence. Chromatograms from each sequence variation are generated by the Sequence Navigator® software and printed on a color printer. The peaks are interpreted by the first primary technologist and a second primary technologist. A secondary technologist then reviews the chromatograms. The results are finally interpreted by a geneticist. In each instance, a variation is compared to known polymorphisms for position and base change. If the sample BRCA1 sequence matches the BRCA1<sup>(omi1)</sup> SEQ. ID. NO.:1 standard, with only variations within the known list of polymorphisms, it is interpreted as a gene sequence.

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### EXAMPLE 3

#### DETERMINING THE ABSENCE OF A MUTATION IN THE BRCA1 GENE USING BRCA1<sup>(omi1)</sup> AND SEVEN POLYMORPHISMS FOR REFERENCE

A person skilled in the art of genetic susceptibility testing will find the present invention useful for determining the presence of a known or previously unknown mutation in the BRCA1 gene. A list of mutations of BRCA1 is publicly available in the Breast Cancer Information Core at:

[http://www.nchgr.nih.gov/dir/lab\\_transfer/bic](http://www.nchgr.nih.gov/dir/lab_transfer/bic). This data site became publicly available on November 1, 1995. Friend, S. *et al. Nature Genetics* 11:238, (1995).

Sequencing is carried out as in EXAMPLE 1 using a blood sample from the patient in question. However, a BRCA1<sup>(omi)</sup> sequence is used for reference and polymorphic sites are compared to the nucleic acid sequences listed above for codons at each polymorphic site. A sample is one which compares to the BRCA1<sup>(omi2)</sup> SEQ. ID. NO.: 3 sequence and contains one of the base variations which occur at each of the polymorphic sites. The codons which occur at each of the polymorphic sites are paired here reference.

30

- AGC and AGT at position 2201,
- TTG and CTG at position 2430,

- CCG and CTG at position 2731,
- GAA and GGA at position 3232,
- AAA and AGA at position 3667,
- TCT and TCC at position 4427, and
- 5 • AGT and GGT at position 4956.

The availability of these polymorphic pairs provides added assurance that one skilled in the art can correctly interpret the polymorphic variations without mistaking a variation for a mutation.

Exon 11 of the BRCA1 gene is subjected to direct dideoxy sequence analysis by  
 10 asymmetric amplification using the polymerase chain reaction (PCR) to generate a single stranded product amplified from this DNA sample. Shuldiner, *et al.*, Handbook of Techniques in Endocrine Research, p. 457-486, DePablo, F., Scanes, C., eds., Academic Press, Inc., 1993. Fluorescent dye is attached for automated sequencing using the Taq Dye Terminator® Kit (Perkin-Elmer cat# 401628). DNA sequencing is performed in  
 15 both forward and reverse directions on an Applied Biosystems, Inc. (ABI) automated Model 377® sequencer. The software used for analysis of the resulting data is "Sequence Navigator® software" purchased through ABI.

#### 1. Polymerase Chain Reaction (PCR) Amplification

20 Genomic DNA (100 nanograms) extracted from white blood cells of the subject is amplified in a final volume of 25 microliters containing 1 microliter (100 nanograms) genomic DNA, 2.5 microliters 10X PCR buffer (100 mM Tris, pH 8.3, 500 mM KCl, 1.2 mM MgCl<sub>2</sub>), 2.5 microliters 10X dNTP mix (2 mM each nucleotide), 2.5 microliters forward primer (BRCA1-11K-F, 10 micromolar solution), 2.5 microliters reverse primer  
 25 (BRCA1-11K-R, 10 micromolar solution), and 1 microliter Taq polymerase (5 units), and 13 microliters of water.

The PCR primers used to amplify a patient's sample BRCA1 gene are listed in Table II. The primers were synthesized on an DNA/RNA Model 394® Synthesizer. Thirty-five cycles of amplification are performed, each consisting of denaturing (95°C; 30  
 30 seconds), annealing (55°C; 1 minute), and extension (72°C; 90 seconds), except during the first cycle in which the denaturing time is increased to 5 minutes, and during the

last cycle in which the extension time is increased to 5 minutes.

5 PCR products are purified using Qia-quick® PCR purification kits (Qiagen, cat# 28104; Chatsworth, CA). Yield and purity of the PCR product determined spectrophotometrically at OD<sub>260</sub> on a Beckman DU 650 spectrophotometer.

## 2. Dideoxy Sequence Analysis

10 Fluorescent dye is attached to PCR products for automated sequencing using the Taq Dye Terminator® Kit (Perkin-Elmer cat# 401628). DNA sequencing is performed in both forward and reverse directions on an Applied Biosystems, Inc. (ABI) Foster City, CA., automated Model 377® sequencer. The software used for analysis of the resulting data is "Sequence Navigator® software" purchased through ABI. The BRCA1<sup>(omi2)</sup> SEQ. ID. NO.: 3 sequence is entered into the Sequence Navigator® software as the Standard for comparison. The Sequence Navigator® software compares the sample sequence to 15 the BRCA1<sup>(omi2)</sup> SEQ. ID. NO.: 3 standard, base by base. The Sequence Navigator® software highlights all differences between the BRCA1<sup>(omi2)</sup> SEQ. ID. NO.: 3 DNA sequence and the patient's sample sequence.

A first technologist checks the computerized results by comparing visually the BRCA1<sup>(omi2)</sup> SEQ. ID. NO.: 3 standard against the patient's sample, and again highlights 20 any differences between the standard and the sample. The first primary technologist then interprets the sequence variations at each position along the sequence. Chromatograms from each sequence variation are generated by the Sequence Navigator® software and printed on a color printer. The peaks are interpreted by the first primary technologist and also by a second primary technologist. A secondary 25 technologist then reviews the chromatograms. The results are finally interpreted by a geneticist. In each instance, a variation is compared to known polymorphisms for position and base change. If the sample BRCA1 sequence matches the BRCA1<sup>(omi2)</sup> SEQ. ID. NO.: 3 standard, with only variations within the known list of polymorphisms, it is interpreted as a gene sequence.

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## EXAMPLE 4

DETERMINING THE PRESENCE OF A MUTATION IN THE BRCA1 GENE USING BRCA1<sup>(oml)</sup> AND SEVEN POLYMORPHISMS FOR REFERENCE

A person skilled in the art of genetic susceptibility testing will find the present invention useful for determining the presence of a known or previously unknown mutation in the BRCA1 gene. A list of mutations of BRCA1 is publicly available in the Breast Cancer Information Core at:

http://www.nchgr.nih.gov/dir/lab\_transfer/bic. This data site became publicly available on November 1, 1995. Friend, S. *et al. Nature Genetics* 11:238, (1995). In this example, a mutation in exon 11 is characterized by amplifying the region of the mutation with a primer which matches the region of the mutation.

Exon 11 of the BRCA1 gene is subjected to direct dideoxy sequence analysis by asymmetric amplification using the polymerase chain reaction (PCR) to generate a single stranded product amplified from this DNA sample. Shuldiner, *et al.*, Handbook of Techniques in Endocrine Research, p. 457-486, DePablo, F., Scanes, C., eds., Academic Press, Inc., 1993. Fluorescent dye is attached for automated sequencing using the Taq Dye Terminator<sup>®</sup> Kit (Perkin-Elmer cat# 401628). DNA sequencing is performed in both forward and reverse directions on an Applied Biosystems, Inc. (ABI) automated Model 377<sup>®</sup> sequencer. The software used for analysis of the resulting data is "Sequence Navigator<sup>®</sup> software" purchased through ABI.

#### 1. Polymerase Chain Reaction (PCR) Amplification

Genomic DNA (100 nanograms) extracted from white blood cells of the subject is amplified in a final volume of 25 microliters containing 1 microliter (100 nanograms) genomic DNA, 2.5 microliters 10X PCR buffer (100 mM Tris, pH 8.3, 500 mM KCl, 1.2 mM MgCl<sub>2</sub>), 2.5 microliters 10X dNTP mix (2 mM each nucleotide), 2.5 microliters forward primer (BRCA1-11K-F, 10 micromolar solution), 2.5 microliters reverse primer (BRCA1-11K-R, 10 micromolar solution), and 1 microliter Taq polymerase (5 units), and 13 microliters of water.

The PCR primers used to amplify segment K of exon 11 (where the mutation is found) are as follows:

BRCA1-11K-F: 5'-GCA AAA GCG TCC AGA AAG GA-3' SEQ ID NO:69

BRCA1-11K-R: 5'-AGT CTT CCA ATT CAC TGC AC-3' SEQ ID NO:70

The primers are synthesized on an DNA/RNA Model 394® Synthesizer.

5 Thirty-five cycles are performed, each consisting of denaturing (95°C; 30 seconds), annealing (55°C; 1 minute), and extension (72°C; 90 seconds), except during the first cycle in which the denaturing time is increased to 5 minutes, and during the last cycle in which the extension time is increased to 5 minutes.

10 PCR products are purified using Qia-quick® PCR purification kits (Qiagen, cat# 28104; Chatsworth, CA). Yield and purity of the PCR product determined spectrophotometrically at OD<sub>260</sub> on a Beckman DU 650 spectrophotometer.

## 2. Dideoxy Sequence Analysis

15 Fluorescent dye is attached to PCR products for automated sequencing using the Taq Dye Terminator® Kit (Perkin-Elmer cat# 401628). DNA sequencing is performed in both forward and reverse directions on an Applied Biosystems, Inc. (ABI) Foster City, CA., automated Model 377® sequencer. The software used for analysis of the resulting data is "Sequence Navigator® software" purchased through ABI. The BRCA1<sup>(omi2)</sup> SEQ. ID. NO.: 3 sequence is entered into the Sequence Navigator® software as the Standard for comparison. The Sequence Navigator® software compares the sample sequence to  
20 the BRCA1<sup>(omi2)</sup> SEQ. ID. NO.: 3 standard, base by base. The Sequence Navigator® software highlights all differences between the BRCA1<sup>(omi2)</sup> SEQ. ID. NO.: 3 DNA sequence and the patient's sample sequence.

25 A first technologist checks the computerized results by comparing visually the BRCA1<sup>(omi2)</sup> SEQ. ID. NO.: 3 standard against the patient's sample, and again highlights any differences between the standard and the sample. The first primary technologist then interprets the sequence variations at each position along the sequence. Chromatograms from each sequence variation are generated by the Sequence Navigator® software and printed on a color printer. The peaks are interpreted by the first primary technologist and a second primary technologist. A secondary technologist  
30 then reviews the chromatograms. The results are finally interpreted by a geneticist. In each instance, a variation is compared to known polymorphisms for position and base

change. Mutations are noted by the length of non-matching variation. Such a lengthy mismatch pattern occurs with deletions and substitutions.

### 3. Result

5 Using the above PCR amplification and standard fluorescent sequencing technology, The 3888delGA mutation may be found. The 3888delGA mutation The BRCA1 gene lies in segment "K" of exon 11. The DNA sequence results demonstrate the presence of a two base pair deletion at nucleotides 3888 and 3889 of the published  
10 BRCA1<sup>(omi)</sup> sequence. This mutation interrupts the reading frame of the BRCA1 transcript, resulting in the appearance of an in-frame terminator (TAG) at codon position 1265. This mutation is, therefore, predicted to result in a truncated, and most likely, non-functional protein. The formal name of the mutation will be 3888delGA. This mutation is named in accordance with the suggested nomenclature for naming mutations, Baudet, A *et al.*, *Human Mutation* 2:245-248, (1993).

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#### EXAMPLE 5

#### USE OF THE BRCA1<sup>(omi1)</sup> GENE THERAPY

The growth of ovarian, breast or prostate cancer can be arrested by increasing the  
20 expression of the BRCA1 gene where inadequate expression of that gene is responsible for hereditary ovarian, breast and prostate cancer. It has been demonstrated that transfection of BRCA1 into cancer cells inhibits their growth and reduces tumorigenesis. Gene therapy is performed on a patient to reduce the size of a tumor. The LXS vector is transformed with any of the BRCA1<sup>(omi1)</sup> SEQ. ID. NO.:1,  
25 BRCA1<sup>(omi2)</sup> SEQ. ID. NO.:3, or BRCA1<sup>(omi3)</sup> SEQ. ID. NO.:5 coding region.

#### Vector

The LXS vector is transformed with wildtype BRCA1<sup>(omi1)</sup> SEQ. ID. NO.:1 coding sequence. The LXS-BRCA1<sup>(omi1)</sup> retroviral expression vector is constructed by cloning  
30 a *SalI*-linked BRCA1<sup>(omi1)</sup> cDNA (nucleotides 1-5711) into the *XhoI* site of the vector LXS. Constructs are confirmed by DNA sequencing. Holt *et al.* *Nature Genetics* 12: 298-302 (1996).

Retroviral vectors are manufactured from viral producer cells using serum free and phenol-red free conditions and tested for sterility, absence of specific pathogens, and absence of replication-competent retrovirus by standard assays. Retrovirus is stored frozen in aliquots which have been tested.

- 5 Patients receive a complete physical exam, blood, and urine tests to determine overall health. They may also have a chest X-ray, electrocardiogram, and appropriate radiologic procedures to assess tumor stage.

Patients with metastatic ovarian cancer are treated with retroviral gene therapy by infusion of recombinant LXS<sub>N</sub>-BRCA1<sup>(omi1)</sup> retroviral vectors into peritoneal sites  
10 containing tumor, between 10<sup>9</sup> and 10<sup>10</sup> viral particles per dose. Blood samples are drawn each day and tested for the presence of retroviral vector by sensitive polymerase chain reaction (PCR)-based assays. The fluid which is removed is analyzed to determine:

- 15 1. The percentage of cancer cells which are taking up the recombinant LXS<sub>N</sub>-BRCA1<sup>(omi1)</sup> retroviral vector combination. Successful transfer of BRCA1 gene into cancer cells is shown by both RT-PCR analysis and *in situ* hybridization.

RT-PCR is performed with by the method of Thompson *et al. Nature Genetics* 2: 444-450 (1995), using primers derived from BRCA1<sup>(omi1)</sup> SEQ. ID. NO.:1. Cell lysates are  
20 prepared and immunoblotting is performed by the method of Jensen *et al. Nature Genetics* 12: 303-308 1996) and Jensen *et al. Biochemistry* 31: 10887-10892 (1992).

2. Presence of programmed cell death using ApoTAG® *in situ* apoptosis detection kit (Oncor, Inc., Gaithersburg, Maryland) and DNA analysis.

- 25 3. Measurement of BRCA I gene expression by slide immunofluorescence or western blot.

Patients with measurable disease are also evaluated for a clinical response to LXS<sub>N</sub>-BRCAI, especially those that do not undergo a palliative intervention immediately after  
30 retroviral vector therapy. Fluid cytology, abdominal girth, CT scans of the abdomen, and local symptoms are followed.



For other sites of disease, conventional response criteria are used as follows:

1. Complete Response (CR), complete disappearance of all measurable lesions and of all signs and symptoms of disease for at least 4 weeks.
2. Partial Response (PR), decrease of at least 50% of the sum of the products of the 2 largest perpendicular diameters of all measurable lesions as determined by 2 observations not less than 4 weeks apart. To be considered a PR, no new lesions should have appeared during this period and none should have increased in size.
3. Stable Disease, less than 25% change in tumor volume from previous evaluations.
4. Progressive Disease, greater than 25% increase in tumor measurements from prior evaluations.

The number of doses depends upon the response to treatment.

For further information related to this gene therapy approach see in "BRCA1 Retroviral Gene Therapy for Ovarian Cancer" a Human Gene Transfer Protocol: NIH ORDA Registration #: 9603-149 Jeffrey Holt, JT, M.D. and Carlos L. Arteaga, M.D.

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“Breast and Ovarian cancer” is understood by those skilled in the art to include breast and ovarian cancer in women and also breast and prostate cancer in men. BRCA1 is associated genetic susceptibility to inherited breast and ovarian cancer in women and also breast and prostate cancer in men. Therefore, claims in this document

15 which recite breast and/or ovarian cancer refer to breast, ovarian and prostate cancers in men and women. Although the invention has been described with reference to the presently preferred embodiments, it should be understood that various modifications can be made without departing from the spirit of the invention. Accordingly, the invention is limited only by the following claims.